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Therapeutic administration of broadly neutralizing FI6 antibody reveals lack of interaction between human IgG1 and pig Fc receptors

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The authors declare a potential conflict of interest and state it below

DC, EC, GA, BG are employees of Humabs Biomed, a company that develops anti-infectives human monoclonal antibodies. All other authors declare no competing interests. Author Contributions

Author contribution statement

ET, SM, JH, BH, DC, BC designed and performed the experiments and analysed the data. EP collected pathology samples and FJS performed the pathological analysis. EC, GA, BG produced the FI6 antibody and provided advice on ADCC. JS provide sequence alignments. AT and PR provided reagents for and crucial advice on microneutralization assays. SM, ET and DC wrote and edited the paper.

Keywords

Influenza1, Anti-stem antibody2, PIG3, Fc receptor4, FI65, Enhanced disease6

Abstract

Word count: 210

Influenza virus infection is a significant global health threat. Because of the lack of cross protective universal vaccines, short time window during which antivirals are effective and drug resistance, new therapeutic anti-influenza strategies are required. Broadly cross-protective antibodies that target conserved sites in the hemagglutinin (HA) stem region, have been proposed as therapeutic agents. FI6 is the first proven such monoclonal antibody to bind to H1-H16 and is protective in mice and ferrets. Multiple studies have shown that Fc-dependent mechanisms are essential for FI6 in vivo efficacy. Here we show that therapeutic administration of FI6 either intravenously or by aerosol to pigs did not reduce viral load in nasal swabs or broncho-alveolar lavage, but aerosol delivery of FI6 reduced gross pathology significantly. We demonstrate that pig Fc receptors do not bind human IgG1 and that FI6 did not mediate antibody dependent cytotoxicity (ADCC) with pig PBMC, confirming that ADCC is an important mechanism of protection by anti-stem antibodies in vivo. Enhanced respiratory disease, which has been associated in pigs with cross-reactive non-neutralising anti-HA antibodies, did not occur after FI6 administration. Our results also show that in vitro neutralizing antibody responses are not a robust correlate of protection for the control of influenza infection and pathology in a natural host model.

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Animal experiments were approved by the Pirbright Institute ethics committee, according to the UK Animal (Scientific Procedures) Act 1986.

In review

Therapeutic administration of broadly neutralizing FI6 antibody reveals lack of interaction between human IgG1 and pig Fc receptors

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Keywords: Influenza¹, Anti-stem antibody², Pig³, Fc receptor⁴, FI6⁵, Enhanced disease⁶.

Abstract

Influenza virus infection is a significant global health threat. Because of the lack of cross protective universal vaccines, short time window during which antivirals are effective and drug resistance, new therapeutic anti-influenza strategies are required. Broadly cross-protective antibodies that target conserved sites in the hemagglutinin (HA) stem region, have been proposed as therapeutic agents. FI6 is the first **proven** such monoclonal antibody **to bind to H1-H16** and is protective in mice and ferrets. Multiple studies have shown that Fc-dependent mechanisms are essential for FI6 *in vivo* efficacy. Here we show that therapeutic administration of FI6 either intravenously or by aerosol to pigs did not reduce viral load in nasal swabs or broncho-alveolar lavage, but aerosol delivery of FI6 reduced gross pathology significantly. We demonstrate that pig Fc receptors do not bind human IgG1 and that FI6 did not mediate antibody dependent cytotoxicity (ADCC) with pig PBMC, confirming that ADCC is an important mechanism of protection by anti-stem antibodies *in vivo*. Enhanced respiratory disease, which has been associated in pigs with cross-reactive non-neutralising anti-HA antibodies, did not occur after FI6 administration. Our results also show that *in vitro* neutralizing antibody responses are not a robust correlate of protection for the control of influenza infection and pathology in a natural host model.

Introduction

Influenza virus infection and immunization induce protective antibody responses. A major part of the antibody response is directed at the hemagglutinin (HA) glycoprotein. Influenza HA is composed of

two domains: the immunodominant globular head, which is strain specific and the stalk which is relatively conserved within each subtype. Seasonal immunization induces antibodies predominantly against the globular head which neutralize the immunizing strain very effectively, but escape variants rapidly emerge and are responsible for antigenic drift. In the last decade influenza-neutralizing antibodies that target conserved sites in the HA stem of influenza A viruses (IAVs) have been described and these show cross-reactivity between group 1 and group 2 viruses (1-7). Anti-stem antibodies are less potent at direct viral neutralization as compared to anti-head antibodies, but they mediate protection *in vivo* through Fc-dependent effector functions, which can be assessed *in vitro* by measuring antibody-dependent cellular cytotoxicity (ADCC), complement-dependent cytotoxicity (CDC) or antibody-dependent cellular phagocytosis (ADCP) (4, 8, 9). FI6 was the first proven broadly neutralizing antibody to be described, capable of recognizing the HAs of all 16 subtypes and neutralizing both group 1 and 2 IAVs (4). Passive transfer of FI6 conferred protection in mice and ferrets. It has been proposed that such broadly cross-reactive antibodies might have potential as therapeutic agents for treatment of severe influenza and several are tested in clinical trials (10, 11).

A potential problem of developing such antibodies as immune therapeutics is enhanced respiratory disease and increased pathology, associated with immune complexes of low avidity or non-neutralizing antibodies. Vaccine-associated enhanced respiratory disease (VAERD) has been observed in pigs when heterologous IAV infection occurs after immunization with mismatched whole inactivated vaccine (WIV) (12-15). VAERD was associated with the presence of high titer cross-reacting non-neutralizing antibodies targeting the conserved stem domain at a site adjacent to the fusion peptide. In the absence of neutralizing antibodies against the globular head of H1N1pdm09, stem antibodies were associated increased virus infection of MDCK cells *in vitro* and enhanced membrane fusion (16).

As both pigs and humans are readily infected with IAVs of similar subtype, the pig is an appropriate model for investigating both swine and human disease. Like humans, pigs are outbred, and physiologically, anatomically and immunologically similar to humans. The porcine lung also resembles the human in terms of its physiology, morphology and distribution of receptors bound by IAV (17, 18). Here we used the pig influenza model to test whether therapeutic administration of FI6 would reduce or enhance disease.

Materials and Methods

Animals and influenza virus challenge

Animal experiments were approved by the Pirbright Institute ethics committee, according to the UK Animal (Scientific Procedures) Act 1986. Five to six week old landrace cross, female pigs were obtained from a commercial high health status herd. Pigs were screened for absence of IAV infection by matrix (M) gene real-time quantitative reverse transcriptase polymerase chain reaction (qRT-PCR) (19) and antibody-free status was confirmed using haemagglutination inhibition (HAI) with 4 swine IAV antigens – pandemic H1N1, H1N2, H3N2 and avian-like H1N1. Pigs weighed between 9 and 12 kg. All pigs were challenged with 1×10^7 plaque forming units (PFU) of A/sw/Eng/1353/09 (pdmH1N1) influenza virus strain. The pigs were inoculated by the intra-nasal route using a mucosal atomization device, MAD300 (Wolfe Tory Medical) with 2ml of virus administered to each nostril. The virus was propagated in Madin-Darby canine kidney (MDCK) cells. The challenged pigs were randomly divided into five groups of 5 animals and received the following antibodies (experimental design in Fig 1A). 1) Control group – no treatment; 2) 15 mg/kg of FI6 antibody intra-venously (FI6 I.V.) in the ear vein at 1 day post infection (dpi); 3) 1.5 mg/ml FI6 antibody administered by aerosol (FI6 aer) using Innospire Mini (Philips Respironics http://evergreen-nebulizers.co.uk/respironics/innospire_mini.html) with Aerogen mesh

reservoir with an airspeed of 2 L/min at 1 and 2 days post infection (dpi); 4) 15 mg/kg of EVB114 antibody I.V. in the ear vein at 1 dpi and 5) 1.5 mg/kg of the MPE8 antibody by aerosol at 1 and 2 dpi as described above. All antibodies were provided by Humabs BioMed. They were produced in Chinese hamster ovary (CHO) cells, affinity-purified using HiTrap Protein A columns (GE Healthcare) followed by desalting using HiTrap Fast desalting columns (GE Healthcare). The final product were sterilized by filtration through 0.22 µm filters and stored at +4°C until use. Antibodies were diluted in phosphate buffered saline (PBS) to the desired concentration before administration. Animals were monitored by observing demeanour, appetite and respiratory signs such as coughing and sneezing.

Gross pathology and histopathological scoring of lung lesions

Animals were humanely killed 4 dpi with an overdose of pentobarbital sodium anaesthetic. At *post mortem* the lungs were removed and digital photographs taken of the dorsal and ventral aspects. Macroscopic pathology scoring was performed blind using Nikon-NIS Br software to determine the proportion of the total surface area of each lung lobe affected by typical influenza-like gross lesions. Five lung tissue samples per animal from the right lung (2 pieces from apical lobe, 1 from the medial, 1 from the diaphragmatic and 1 from the accessory) were collected into 10% neutral buffered formalin for routine histological processing at the University of Surrey. Formalin fixed tissues were paraffin wax-embedded and 4-µm sections were cut and routinely stained with haematoxylin and eosin. Histopathological changes in the stained lung tissue sections were scored by a veterinary pathologist blinded to the treatment group. Lung histopathology was scored using five parameters (necrosis of the bronchiolar epithelium, airway inflammation, perivascular/bronchiolar cuffing, alveolar exudates and septal inflammation) scored on a 5-point scale of 0 to 4 and then summed to give a total slide score ranging from 0 to 20 and a total animal score from 0 to 100. Scoring criteria were based upon a previously published method (20).

Tissue sample processing

Four nasal swabs (NS) (two per nostril) were taken at 0, 1, 2, 3, 4 dpi. The swabs were placed into 2 ml of virus transport medium comprising tissue culture medium 199 (Sigma-Aldrich) supplemented with 25mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES), 0.035% sodium bicarbonate, 0.5% bovine serum albumin (BSA), penicillin 100 IU/ml, streptomycin 100 µg/ml and nystatin 0.25 µg/ml, vortexed, centrifuged to remove debris and stored at -80°C for subsequent virus titration. Serum samples were collected at the start of the study (prior to challenge) and at 2 and 4 dpi. For Fc binding and ADCC assays blood from healthy humans and uninfected pigs was used. Heparinized blood samples were diluted 1:1 in PBS before density gradient centrifugation. PBMC were harvested from the interface, washed and red blood cells lysed with ammonium chloride lysis buffer, washed again and used in Fc binding and ADCC assays described below. Broncho-alveolar lavage (BAL) was collected from the entire left lung with 150ml of virus transport medium (described above). BAL samples were centrifuged at 300 x g for 15 minutes, supernatant was removed, aliquoted and frozen for antibody analysis.

Virus titration

Viral titers in nasal swabs and BAL were determined by plaque assay on MDCK cells. Duplicate samples were 10-fold serially diluted in Dulbecco's modified Eagles medium (DMEM) and 100 µl of each dilution added to confluent MDCK cells in 12 well tissue culture plates. After 1 hour, the plates were washed and overlaid with 2 ml 1:3 2% (w/v) agarose:medium. Plates were incubated at 37°C for 48 hours, plaques visualized by staining the monolayer with 0.1% (v/v) crystal violet and enumerated. RNA was extracted using the QIAamp viral RNA mini kit (Qiagen) according to the manufacturer's protocol

and viral titers in nasal swabs and BAL fluid was also determined by real-time qRT-PCR amplification of the M gene using PCR conditions as previously described (21). Forward primer sequence AGA TGA GTC TTC TAA CCG AGG TCG, reverse primer sequence TGC AAA GAC ACT TTC CAG TCT CTG and probe sequence FAM-TCA GGC CCC CTC AAA GCC GA-TAMRA

Enzyme-linked immunosorbent assay (ELISA)

Human IgG1 antibody levels in serum and BAL fluid were determined by IgG1 Ready-SET-Go! ELISA (Affymetrix, eBioscience) according to the manufacturer's instructions. After heat inactivation (56°C for 30 min) samples were diluted 1:40 (serum) and 1:2 (BAL fluid). Influenza-specific human Ab titers in serum and BAL fluid were determined by ELISA as previously described (20) with the following modifications. The IgG ELISA was performed in 96-well ELISA plates (BD Biosciences) coated with 1×10^6 PFU/ml of A/swine/England/1353/09 over night at 4°C. Two-fold dilutions of BAL fluid samples or serum (heat inactivated for 30 min at 56°C) were added, starting from 1:2 or 1:10 dilution respectively. Binding of influenza specific Abs was detected using a monoclonal anti-human IgG (Fc) (Biorad) and 3,3',5,5'-tetramethylbenzidine (TMB) substrate (BioLegend). Optical density (OD) readings were taken at 450 and 570 nm (wavelength correction). Ab values were expressed as endpoint titers defined as the highest dilution at which the OD was higher than twice the background OD.

Fc binding

To determine if FI6 was able to bind pig Fc receptors FI6, MPE8 and serum from influenza negative and immune (14 dpi) animals were incubated at 37°C for 1 hour with and without influenza virus. Human and pig PBMC were added and incubated for a further hour at 4°C. Human PBMC were stained with near-infrared fixable Live/Dead (Invitrogen) and anti-human IgG AF488 (HP6017, Biolegend) for 20 minutes at 4°C. Pig PBMC were stained with near-infrared fixable Live/Dead (Invitrogen), CD3 AF647 (BB23-8E6-8C8, BD), CD8 α Pe (76-2-11, BD) anti-human IgG or anti-pig IgG FITC (BIO-RAD). Samples were run on a BD LSR Fortessa and data analysed using FlowJo (Treestar).

Entry microneutralization assay

Serum and BAL fluid were heat inactivated at 56°C for 30 minutes, serially diluted 1:2 in 50 μ l PBS, starting dilution 1:40 for serum and 1:4 for BALF, before addition of 50 μ l green-fluorescent protein (GFP) - H1 virus diluted in virus growth medium (22). Following incubation for 2 hours at 37°C 3×10^4 indicator MDCK-SIAT1 cells were added in a volume of 100 μ l virus growth medium without trypsin and incubated overnight at 37°C. Plates were fixed using 4% paraformaldehyde and GFP fluorescence intensity (FI) was measured at an excitation of 483 nm and an emission of 515 nm. Serum and BALF from animals 14 days post influenza challenge and purified FI6 antibody were included as positive controls.

Antibody-dependent cell-mediated cytotoxicity (ADCC) assay

MDCK- 2,6-sialtransferase (SIAT1) stably transduced with the lentiviral vector pHR-SIN engineered to express the full-length open-reading frame (ORF) of HA from A/Eng/195/2009 were used as target cells for the ADCC assay (22). The HA from A/Eng/195/2009 differs by a single exposed residue at D222G from the Eng/1353 that was used to challenge the pigs. MDCK-HA cells were seeded in round-bottom 96 well plates and incubated with different dilutions of heat-inactivated serum (1:10, 1:20, 1:40, 1:80 or 1:160) or with different amounts of antibody (FI6 or MPE8) for 10 min at 37°C. After that freshly-isolated human or pig PBMCs from healthy donors or animals respectively, were added in a 20 to 1 E:T ratio to the 96 well plates and incubated for 4h at 37°C. MDCK-HA and PBMC were cultured in serum-free

AIM-V medium (Life Technologies, UK). At the end of the incubation period 100 μ l of cell-free supernatant was transferred into a flat-bottom 96 well plate and the Lactate dehydrogenase (LDH) release measured with the Cytotoxicity Detection Kit from Roche according to the manufacturer's instructions. The absorbance was measured at 490 nm and 620 nm in a plate reader. When the purified antibodies MPE8 and FI6 were used the percentage of cytotoxicity on the Y-axis was calculated with the formula: [Sample at each immune antibody dilution with target cells and PBMC minus control antibody at the same dilution with target cells and PBMC] divided by [maximum release of target cells and PBMC in the presence of detergent minus control target and effector spontaneous release without antibody] x 100. In assessing the ADCC activity of serum samples, the percentage of cytotoxicity was calculated as described above but using the naïve sera (corresponding dilution to sera of immunized pigs to calculate the spontaneous release).

Statistical analysis

One-way non-parametric ANOVA (Kruskall-Wallis) with Dunn's post-test for multiple comparisons was performed using GraphPad Prism 6.

Results

Lung pathology and viral load after antibody administration

In order to determine the therapeutic effect of FI6 antibody in the pig influenza model, FI6 was administered I.V. at 15 mg/kg 1 day post infection (dpi). The ebola virus specific antibody, EVB114 was used as a control and delivered at the same concentration I.V. (23). We also administered FI6 as an aerosol (aer) as this route of delivery is highly efficient in targeting the respiratory tract, which is the site of entry and infection of IAV (20, 24-28). We administered 10 times less FI6 by aerosol (1.5 mg/kg) at 1 and 2 dpi. As a control for the aerosol delivery we used MPE8, which is a broadly-neutralizing antibody for human respiratory syncytial virus (HRSV), human metapneumovirus (HMPV), bovine RSV (BRSV) and pneumonia virus of mice (PVM) but not IAV (29) (**Fig. 1A**). All of the mAbs were monoclonal, fully human IgG1. The clinical signs observed were mild and none of the pigs developed moderate or severe disease. The control group showed the most severe gross and histopathology (**Fig. 1B**). A reduction in the gross and histopathology score was observed in all the mAb treated groups. However, this reduction was statistically significant only in gross pathology for the FI6 aer group. Interestingly despite the reduced lung pathology, there were no differences in viral load in nasal swabs at 1, 2, 3 and 4 dpi (**Fig. 2A**) or in the broncho-alveolar lavage (BAL) at the time of sacrifice at 4 dpi (**Fig. 2B**) as determined by plaque forming assays and PCR. This is in contrast with previous studies in mice and ferrets where FI6 administration significantly reduced viral load in the lungs (4).

These results indicate that administration of therapeutic FI6 to pigs did not reduce viral load in NS or BAL but also it did not exacerbate disease as previously shown with anti-stem antibody (16). The mAb treated groups showed reduced pathology, although the reduction was significant only for the aer FI6 group. The reduced pathology is also observed using control antibodies, a finding that might be related to the anti-inflammatory and immunomodulatory activities of human IgG1 Fc. Indeed, it has been shown that the anti-inflammatory activity of human intravenous immunoglobulin is dependent on sialylation of the N-linked glycan of the IgG1 Fc fragment (30, 31).

Influenza binding and neutralizing activity of administered antibodies

ELISA for hum IgG1 confirmed that all antibodies were delivered successfully, albeit the control EVB114 was detected at a lower concentration in the serum perhaps due to differences in the catabolic rates of this mAb. The mAb concentrations declined at 4 dpi compared to 2 dpi, but were still ~107 µg/ml for FI6 and ~45 µg/ml for EVB114 (**Fig. 3A**). Aerosol administration of FI6 and MPE8 did not result in detectable quantities of mAbs in the serum. However, mAbs were detected in BAL, with ~6.5 µg/ml for FI6 and 0.5 µg/ml for MPE8 measured at 4 dpi, 2 days after the last aerosol administration, most likely indicating that the mAbs are catabolized rapidly after aerosol delivery (**Fig. 3A**). Furthermore I.V. FI6 delivery resulted in the presence of ~0.33 µg/ml in the BAL 4 days after the administration of the antibody, approximately 20 fold less as compared to the aer FI6 group. To further confirm the presence and specificity of FI6 after delivery, virus specific ELISA was performed with the challenge virus. As expected influenza specific human mAb was detected in serum after FI6 I.V. delivery at both 2 and 4 dpi, while in BAL a higher titer was seen after aerosol (1:84) compared to FI6 I.V. administration (1:24) (**Fig. 3B**).

To determine whether the FI6 in the serum and BAL was still able to neutralize the virus, which might explain the lack of reduction of viral titer, we performed **entry** virus neutralization using MDCK cells transfected with H1 HA. The serum from the FI6 I.V. group was neutralizing at both 2 and 4 dpi with a mean 50% inhibitory titer of 1:812 at 2 dpi and 1:448 at 4 dpi (**Fig. 4A and B**), comparable to control immune pig serum. In the BAL of the FI6 aer group the mean 50% inhibitory titer was 1:10 at 4 dpi and in the FI6 I.V. group 1:3.4 (although only 2 out of the 5 animals had positive results) (**Fig 4A and C**). The neutralization values for the BAL were lower than a control BAL fluid (1:640) from a pig sacrificed 14 days post challenge with the same virus. No neutralization was detected in the animals receiving control antibodies or in the untreated controls. **Pre-challenge sera from FI6 treated animals and BAL from control animals did not show any neutralizing activity (Fig. 4D).**

Overall these results indicate that the mAbs were successfully delivered and retained their influenza binding and neutralizing activity as measured *in vitro*.

Fc binding and ADCC

As it has been shown convincingly that most broadly neutralizing anti-IAV mAbs mediate their *in vivo* effect through antibody effector functions (4, 9, 32) we next asked whether the FI6 or human IgG1 can bind pig FcR and mediate ADCC. Fc binding was assessed after pre-incubating the mAbs with pdmH1N1 virus in order to form immune complexes. As expected human lymphocytes bound FI6 with 74% of the lymphocytes stained compared to less than 2% for the controls (**Fig. 5A**). In contrast, minimal binding of FI6/pdmH1N1 to pig PBMC was detected. A more detailed analysis was performed by gating on pig NK cells, defined as CD3⁺CD8⁺ (**Fig. 5B and C**), which bound immune pig serum pre-incubated with pdmH1N1, but bound very little FI6 (51.8% for immune pig serum versus 2.37% for FI6). Similar results were obtained after detection of immune complexes with secondary anti-human IgG, indicating that this antibody could bind the pig Ig (**Fig. 5C**).

Finally, to determine whether FI6 can mediate ADCC in pigs we evaluated killing by LDH release from MDCK cells stably transfected with H1 HA. As previously described FI6 was able to mediate ADCC with human PBMC as effector cells (**Fig. 6**), but not with pig PBMC. Immune pig serum from influenza infected or immunized pigs gave specific killing.

These results suggest that the failure of FI6 to protect against influenza infection in pigs is most likely due to the inability of FI6 to bind pig FcR and mediate ADCC, and possibly other effector functions (e.g. ADCP).

248 *Fc binding sites in human, mouse, and pig FcγRs and IgG subclasses*

249 The apparent failure of FI6 to interact functionally with pig FcR led us to compare the putative binding
 250 sites on both the FcγRs and the Fc portion of IgG. Importantly, crystallographic analyses of human IgG
 251 (hIgG) complexed with human FcγRIII and the structure for human FcγRI have elucidated the
 252 important contact sites for this interaction (33-36). On the receptor, the Fc contact sites are spread
 253 across the second immunoglobulin domain, and most notably in the BC, C'E, and FG loops (**Fig. 7A**).
 254 Comparison of known mouse, human, and pig FcγR sequences revealed species-specific variation
 255 within these regions, and does not immediately suggest that mouse FcγRs would have greater affinity
 256 for hIgG than the pig (**Fig. 7A**). However, it has been shown that human IgG1 binds to mouse FcγRIV
 257 and effectively induces ADCC and ADCP with mouse NK cells, mouse polymorphonuclear
 258 leukocytes, and mouse macrophages (37).

259 Among the hIgG subclasses, hIgG1 and hIgG3 bind most strongly to FcγRs; whereas, hIgG2 and hIgG4
 260 bind either poorly or not at all (38). Notably on the Fc portion of the antibody, the lower hinge region,
 261 the hinge-proximal portion of the CH2 domain, and an N-linked glycosylation site in CH2 are
 262 implicated in Fc binding. In particular, the motif "LLGG" in the lower hinge is believed to play a
 263 crucial role (39-41). Indeed, mutation of this motif in hIgG1 to either "VLGG" or "LAGG" was
 264 previously shown to reduce or prevent binding to FcγRI, respectively (42). In addition, the mutation of
 265 residues L234 and L235 to alanine to generate the so called LALA mutant abrogates binding to all FcR
 266 and C1q (43). Conversely, mutation of hIgG2 from "VAG" to "LLGG" and of hIgG4 from "FLGG"
 267 to "LLGG" restored FcγRI binding to levels comparable to hIgG1 (42). Interestingly, mice have fully
 268 conserved this motif in mIgG2a, and have similar motifs in mIgG2b and mIgG3 ("LEGG" and "ILGG",
 269 respectively) (**Fig. 7B**). Of the porcine IgGs, however, pIgG3 is most similar ("VLGA"), whereas the
 270 rest of the subclasses lack this motif, and are generally more similar to hIgG2 in this region. Thus, the
 271 presence of the canonical ("LLGG") FcγR binding motif in both human and mouse IgG, but not in pig
 272 IgG, suggests that porcine FcγRs recognise the Fc portion of IgG differently than in humans and mice.
 273 Structural analysis revealed that the LLGG motif of human IgG1 interacts with hydrophobic residues
 274 (LVG) in the FG loop of the human FcγRIIIB. Similarly, hydrophobic residues are found in human
 275 FcγRIIIB and mouse FcγRIV (LFG and LIG, respectively). Conversely, the pig FcγRIII carries the IIK
 276 motif in the FG loop. The analysis of the interaction of the Fc of human IgG1 with human FcγRIIIB
 277 indicates that the presence of a lysin at position 159, as found in the porcine FcγRIIIB, would clash
 278 with L235, thus interfering with the favorable interaction of the LLGG motif of human IgG1 with the
 279 FG loop required for FcγRIII binding (**Fig. 7C**). This observation might explain the lack of binding of
 280 human IgG1 FI6 to pig PBMCs. Of note, the hinge regions of all pig IgGs (except for pig IGHG3) are
 281 shorter as compared to human IgG1, a finding that might suggest a different modality of interaction of
 282 the pig Fcs with the cognate FG loop of porcine FcγRIII.

283 **Discussion**

284 Our data shows that therapeutic administration of the broadly neutralizing FI6 antibody either
 285 intravenously or by aerosol to pigs did not result in exacerbation of disease. Aerosol delivery of FI6
 286 was the only treatment to reduce gross pathology significantly, although viral titers in nasals swabs or
 287 BAL were unchanged. We further demonstrated that the pig Fc receptors do not bind human IgG1 and
 288 that FI6 did not mediate ADCC with pig PBMC, suggesting that the pig is an inappropriate model to
 289 evaluate human IgG1 antibodies.

Previous studies have shown that all neutralizing and non-neutralizing anti-HA (and anti-neuraminidase) mAbs that recognise a breadth of influenza strains require FcγRs for protection *in vivo* (4, 9, 32), while strain specific mAbs did not. This suggests that the *in vitro* neutralization mechanism of broadly neutralizing mAbs such as inhibition of viral fusion or egress, do not dominate *in vivo* at the doses tested. Our results clearly demonstrate therefore that *in vitro* neutralising antibody responses are not a robust correlate of protection for the control of influenza virus infection and pathology in a natural host model.

There are limited studies describing porcine FcγRs. Although there is obvious overall similarity to their human and mouse counterparts, some FcR in domestic animals are unusual, perhaps most notably bovine Fcγ2R, which although related to other mammalian FcγRs, belongs to a novel gene family and porcine FcγRIIIA, which associates with a molecule that contains significant homology to the cathelin family of antimicrobial proteins (44, 45). Furthermore, the conservation of FcγR binding sites in human and mouse IgG, but not in pig IgG, is consistent with our findings. Clearly differences in interaction with IgG subclasses, cell type and tissue specific expression, as well as species differences should be considered when using these models for *in vivo* evaluation of therapeutic mAbs. Substituting the human Fc portion of the FI6 antibody with a pig Fc would provide definitive proof of the importance of Fc binding and ADCC for therapeutic efficacy of FI6.

It is clear that the delivery of FI6 did not cause pathology or exacerbation of disease as described by Khurana et al (16). In their study the pigs were immunized with a whole irradiation inactivated, adjuvanted H1N2 (human like virus) and challenged with a different pdmH1N109 strain. The authors suggested that the vaccine-induced anti-HA stem antibodies facilitated a conformational change in HA that enhanced its fusion and increased virus entry into cells *in vitro*. Nevertheless because FI6 does not engage FcR mediated effector mechanisms in pigs, it is still possible that these might contribute to VAERD, for example by massive killing of infected cells, leading to inflammation and pathology.

In summary our data shows that therapeutic administration of FI6 or a control, either intravenously or by aerosol to pigs did not exacerbate disease. Aerosol delivery is an effective means of administration for therapeutic mAbs in large animals and possibly humans. FI6 does not bind to pig Fc receptors or mediate ADCC, confirming previous evidence that ADCC is an important mechanism for protection by anti-stem Ab *in vivo*.

Conflict of Interest

DC, EC, GA, BG are employees of Humabs Biomed, a company that develops anti-infectives human monoclonal antibodies. All other authors declare no competing interests. Author Contributions

Author Contributions

ET, SM, JH, BH, DC, BC designed and performed the experiments and analysed the data. EP collected pathology samples and FJS performed the pathological analysis. EC, GA, BG produced the FI6 antibody and provided advice on ADCC. JS provide sequence alignments. AT and PR provided reagents for and crucial advice on microneutralization assays. SM, ET and DC wrote and edited the paper.

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Acknowledgments

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495 Data Availability Statement

496 All datasets for this study are included in the manuscript and the supplementary
497 files.

498 Figure Legends

499 **Fig. 1. Experimental design and lung pathology.** Pigs were infected with A/sw/Eng/1353/09 and
500 received the indicated antibodies either by the I.V. route at 1 dpi or by aerosol at 1 and 2 dpi (A). The
501 animals were sacrificed at 4 dpi and lungs scored for appearance of gross (B) and histopathological
502 lesions (C). Each data point represents an individual within the indicated group and lines represent the
503 mean. * denotes significant difference from the control group ($P < 0.05$). Representative gross
504 pathology of a pig from control group (D) and FI6 aer (E).

505 **Fig. 2. Viral load in nasal washes and BAL.** Pigs were infected with A/sw/Eng/1353/09 and
506 administered the indicated antibodies either by the I.V. route (I.V. solid symbols) or by aerosol (aer,
507 hollow symbols). Nasal swabs (NS) were taken at 0, 1, 2 and 3 dpi and pigs sacrificed at 4 dpi. Viral
508 titers in the nasal swabs (A) and BAL (B) were determined by plaque forming assay (PFU) or real-
509 time qRT-PCR. Each data point represents an individual within the indicated group and bars represent
510 the mean.

511 **Fig. 3. Mucosal and systemic IgG responses following administration of antibodies.** Human IgG1
512 (A) and pdmH1N1 specific IgG (B) titers in serum at 0, 2 and 4 dpi and in BAL at 4 dpi. Note for
513 pdmH1N1 specific antibody in serum – all five pigs had titers of 1:640 at 4dpi, while 2 animals at
514 1:1024 and three at 1:640 at 2dpi.

515 **Fig. 4. Entry neutralization activity of serum and BAL following administration of antibodies.** A)
516 Individual 50% inhibition titers in the serum at 0, 2 and 4 dpi and BAL at 4 dpi. B) Neutralizing
517 antibody response measured in the serum of the pigs following FI6 I.V. administration at 2 and 4 dpi.
518 C) Antibody response in the BAL of animals given FI6 I.V. or by aerosol at 4dpi. D) Negative sera
519 from pre-challenge samples and negative BAL from control animals are shown alongside positive
520 control. The dashed line represents the 50% inhibition titer and FI the fluorescence intensity of GFP.
521 The neutralizing titer of serum and BAL from animals infected with the same A/sw/Eng/1353/09 virus
522 and sacrificed at 14 dpi is shown in red.

523 **Fig. 5: FI6 binding to human and pig Fc receptors.** Antibody and pdmH1N1 virus were pre-
524 incubated for 1 hour at 37°C and then added to the either human or pig PBMC. A) Gated on live cells,
525 singlets and SSCA vs IgG FITC. B) and C) Gated on live cells, singlets and CD3⁺ CD8α⁺ IgG FITC.

526 **Fig. 6: ADCC activity of pig and human PBMC.** MDCK cells expressing H1 HA were incubated
527 with FI6, MPE8, immune or normal pig sera in the presence of either pig or human PBMC. ADCC was
528 measured in triplicate by LDH release. FI6 and MPE8 mAbs were used at 10 µg/ml (left panel) or at a
529 concentration range from 0.1 to 1000 ng/ml (right panel). Representative of three experiments.

530 **Fig. 7. Putative amino acid sequence alignment of FcγR and IgG subclasses.** A) Second IgG
531 domain of human, mouse, and pig Fcγ receptors. B) Lower hinge and CH2 domain of human, mouse,
532 and pig IgG subclasses. Previously reported Fc-FcγR contact sites are shaded. Beta-strands are labelled
533 and shown at top. C) Left. Model of the interaction of the Fc region of human IgG1 (green) with human
534 FcγRIIIb (light blue) (pdb, 1t83). Fc-bound glycans are shown as orange spheres. The L234 and L235
535 Fc residues (purple) and the G159 residue of the FcγRIIIb FG loop are identified. Right. Modelling of
536 the positioning of the porcine K159 residue in the FG loop and its clash with the Fc residue L235

Figure 1.TIF

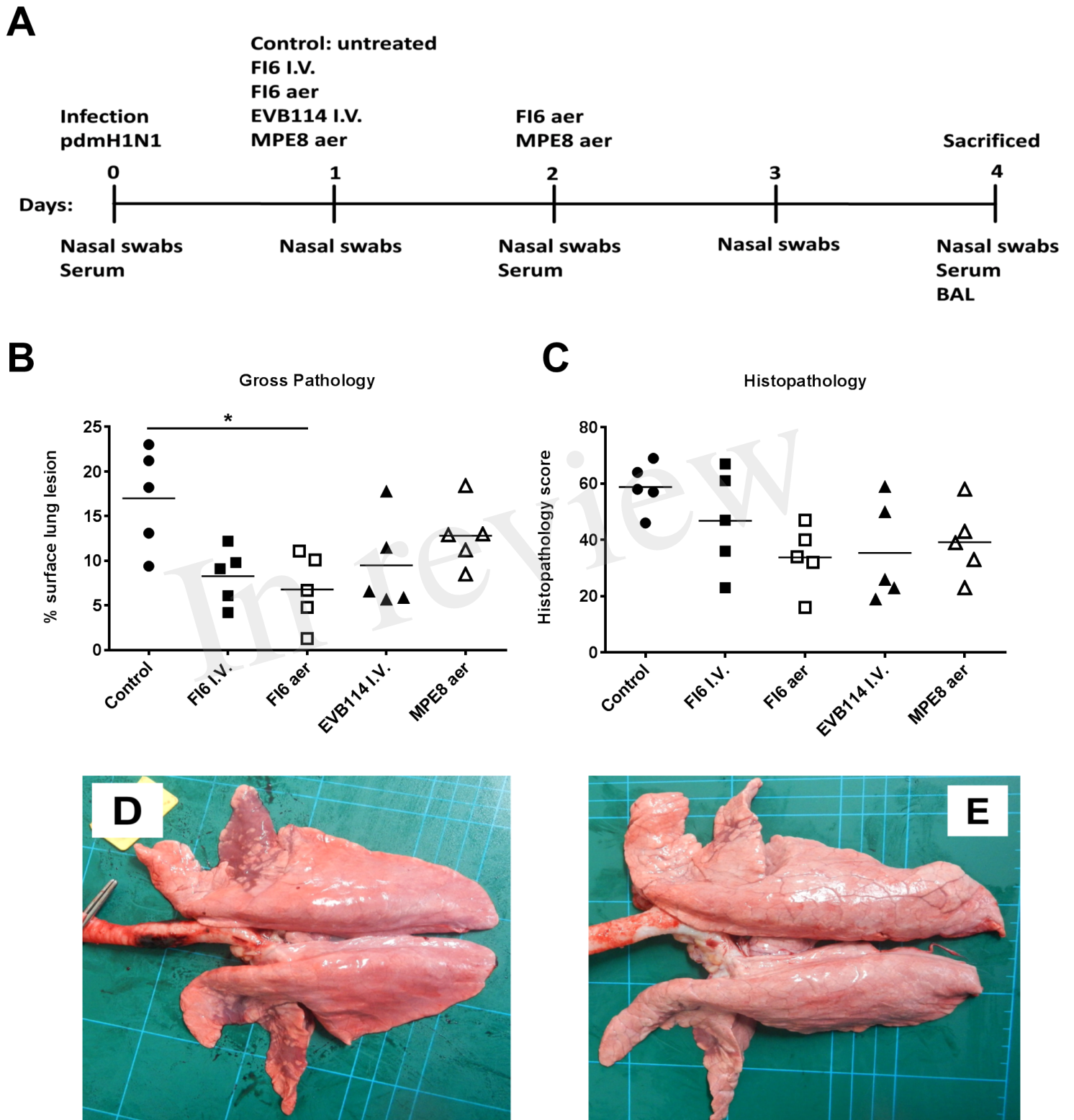


Figure 1

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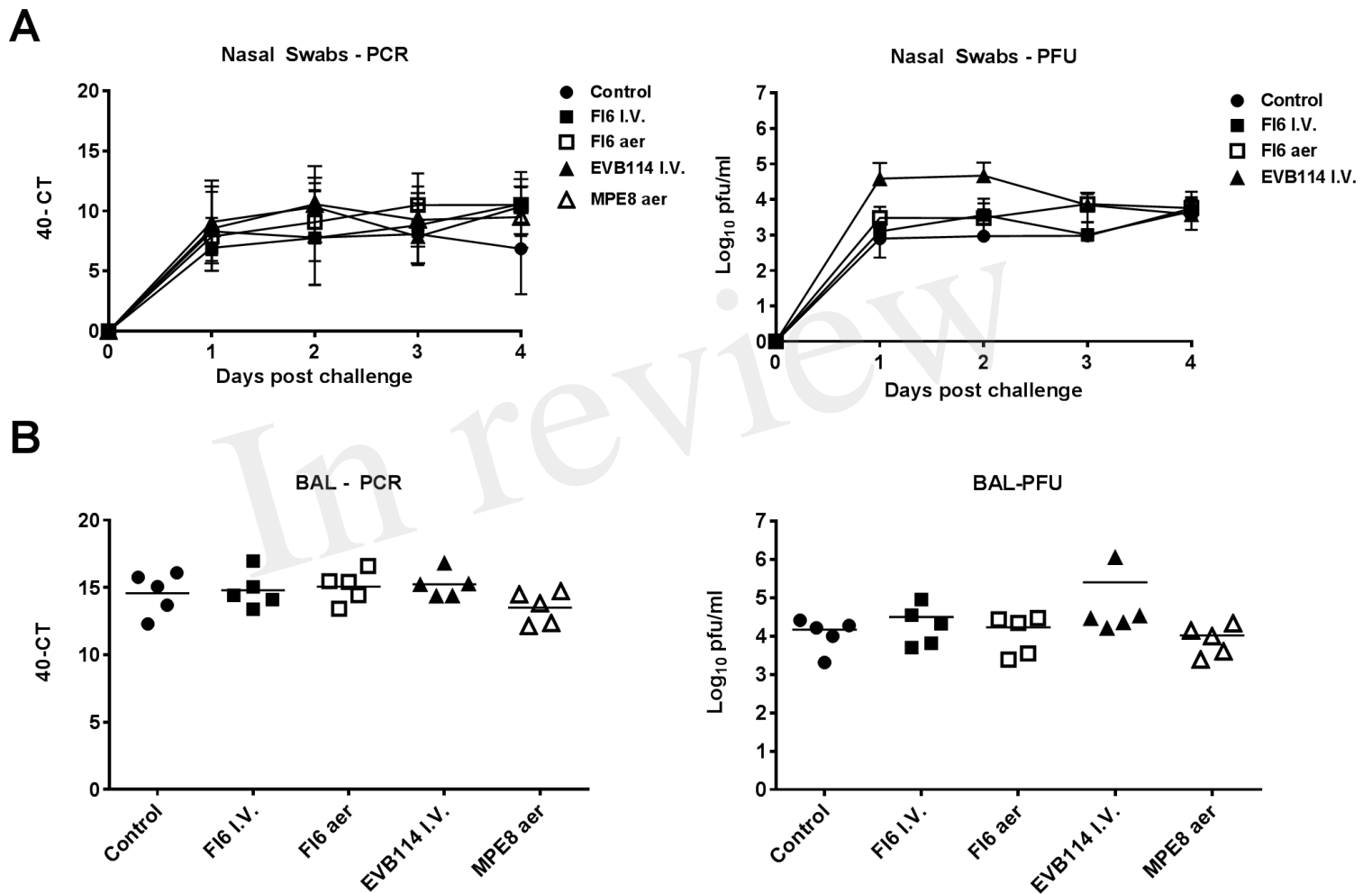


Figure 2

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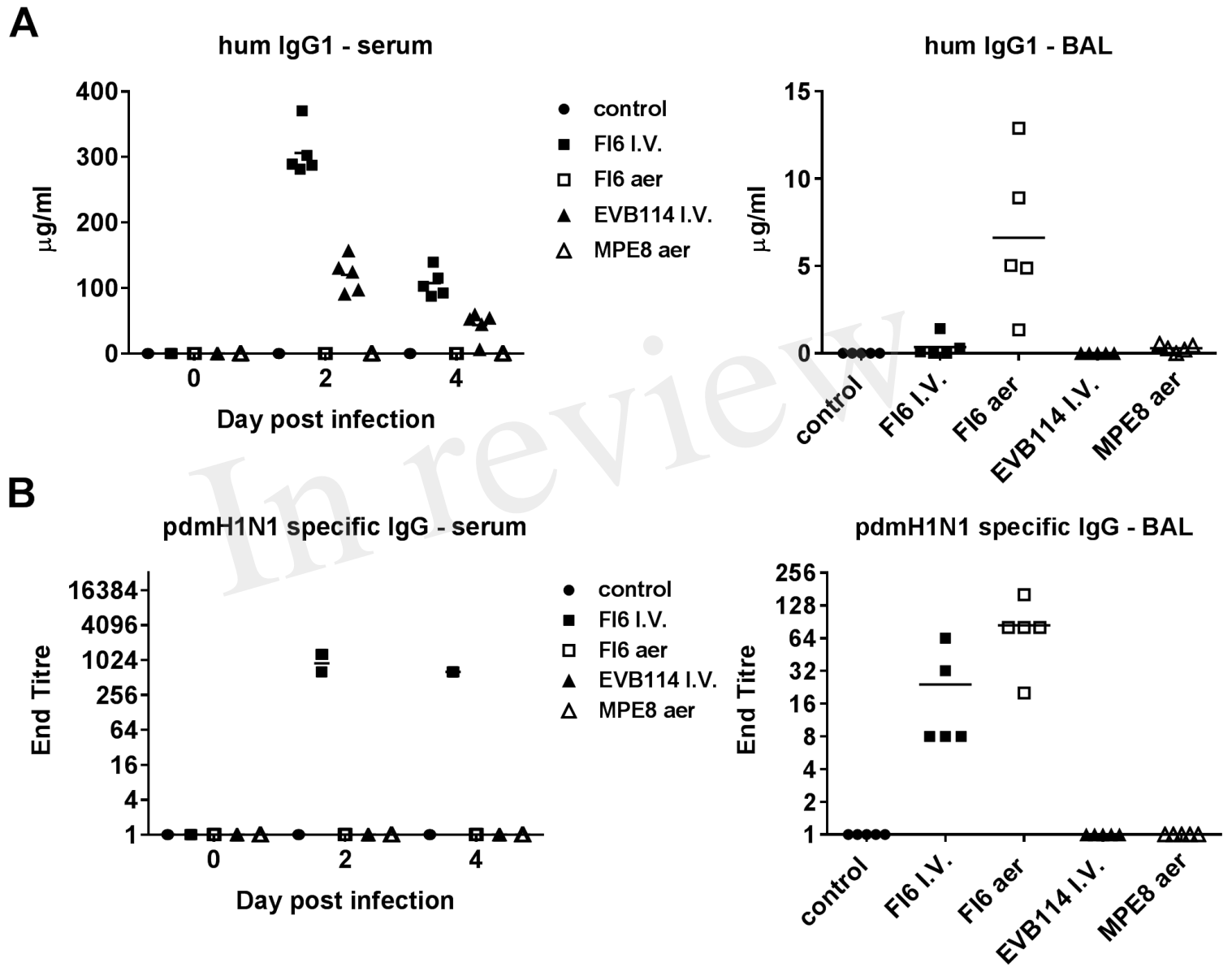


Figure 3

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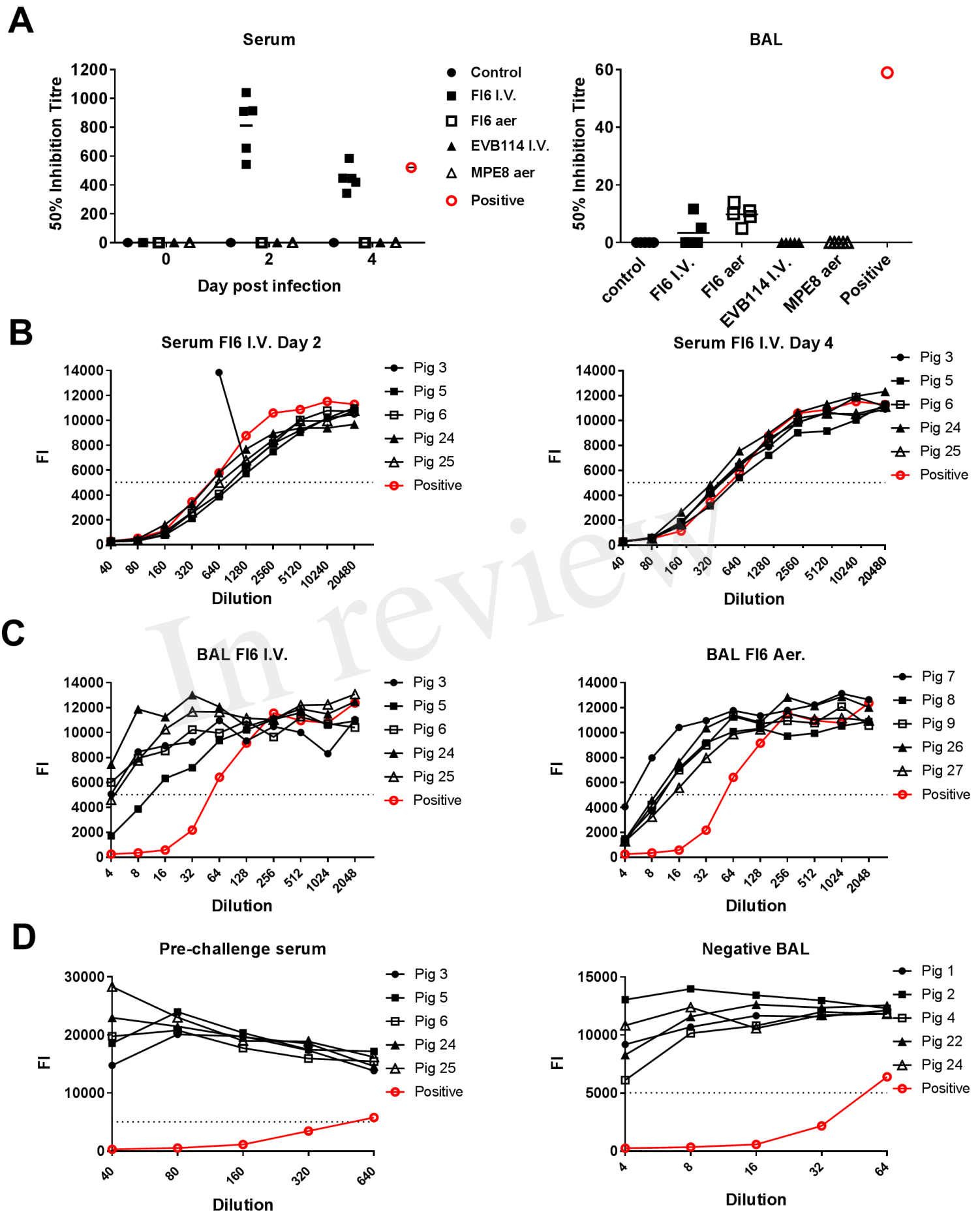
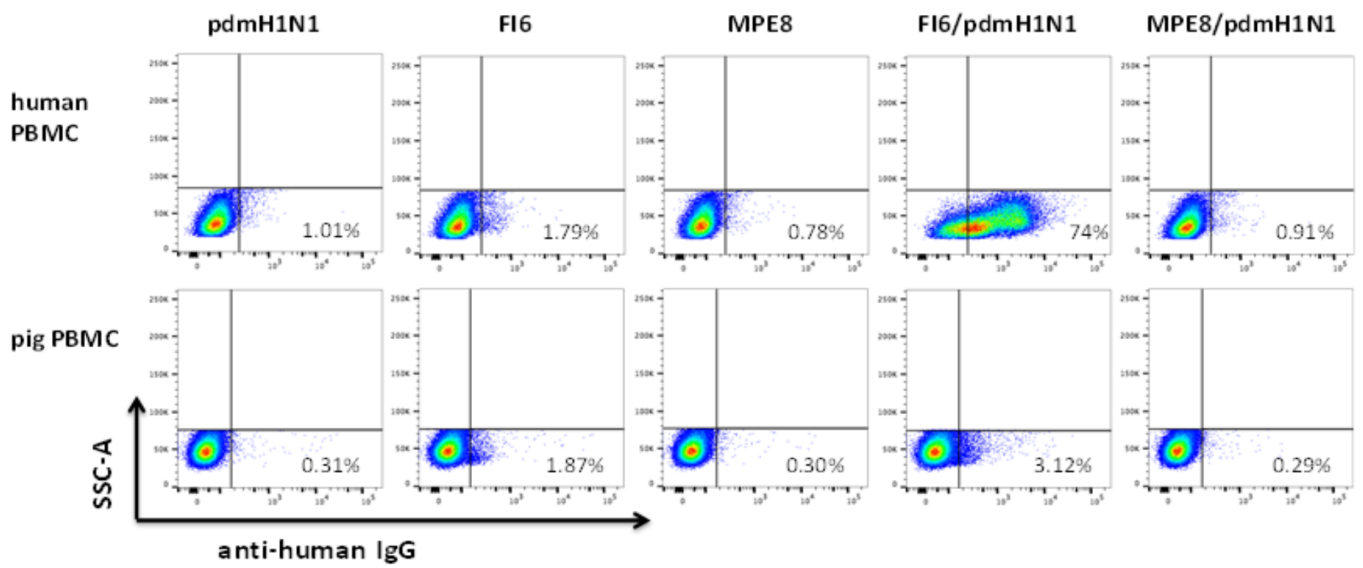


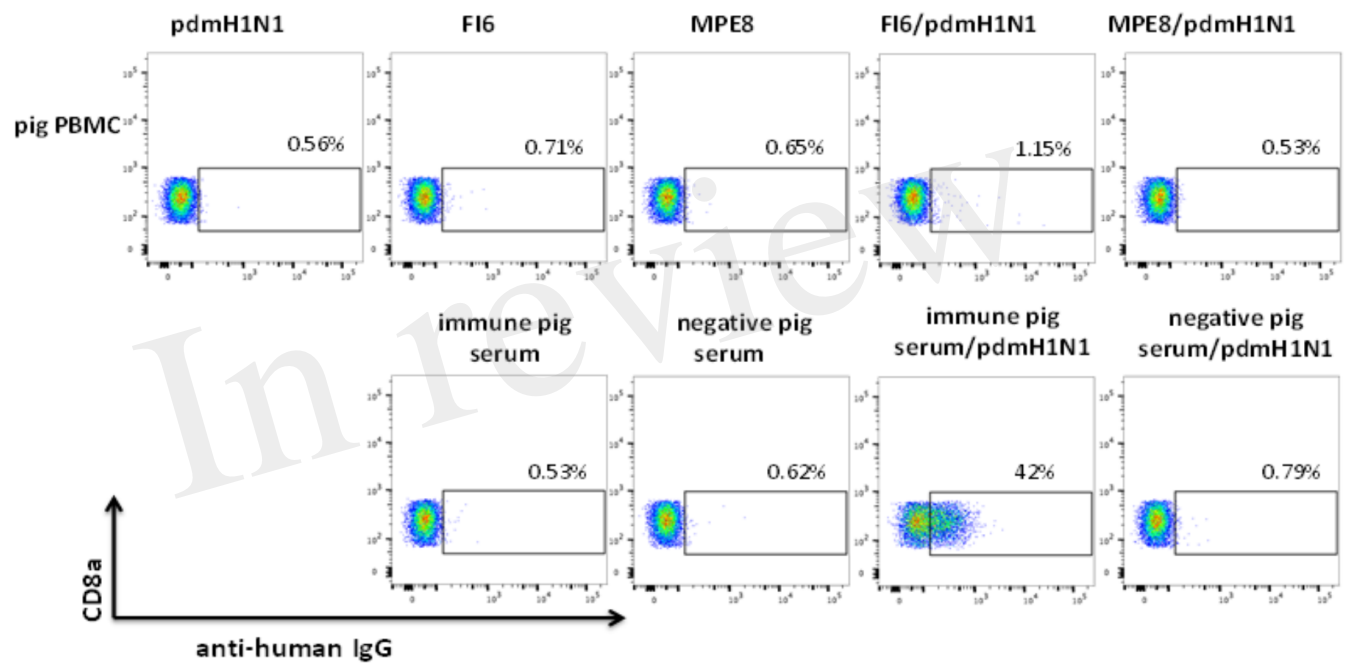
Figure 4

Figure 5.TIF

A



B



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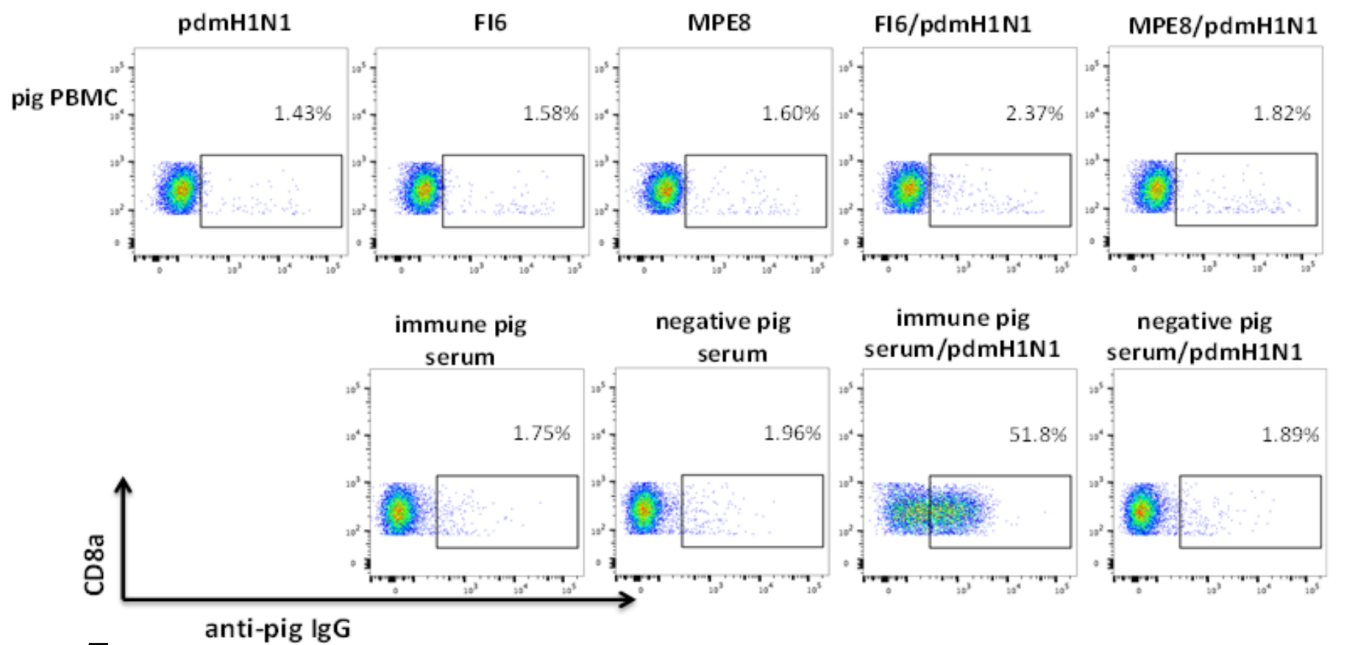


Figure 5

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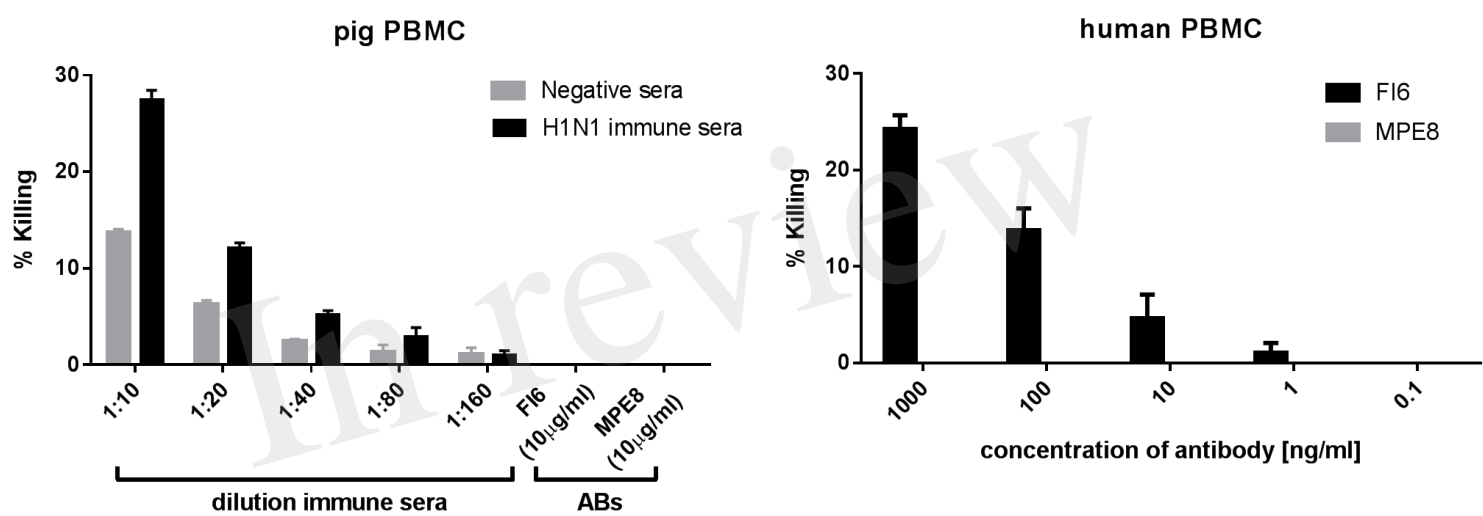


Figure 6

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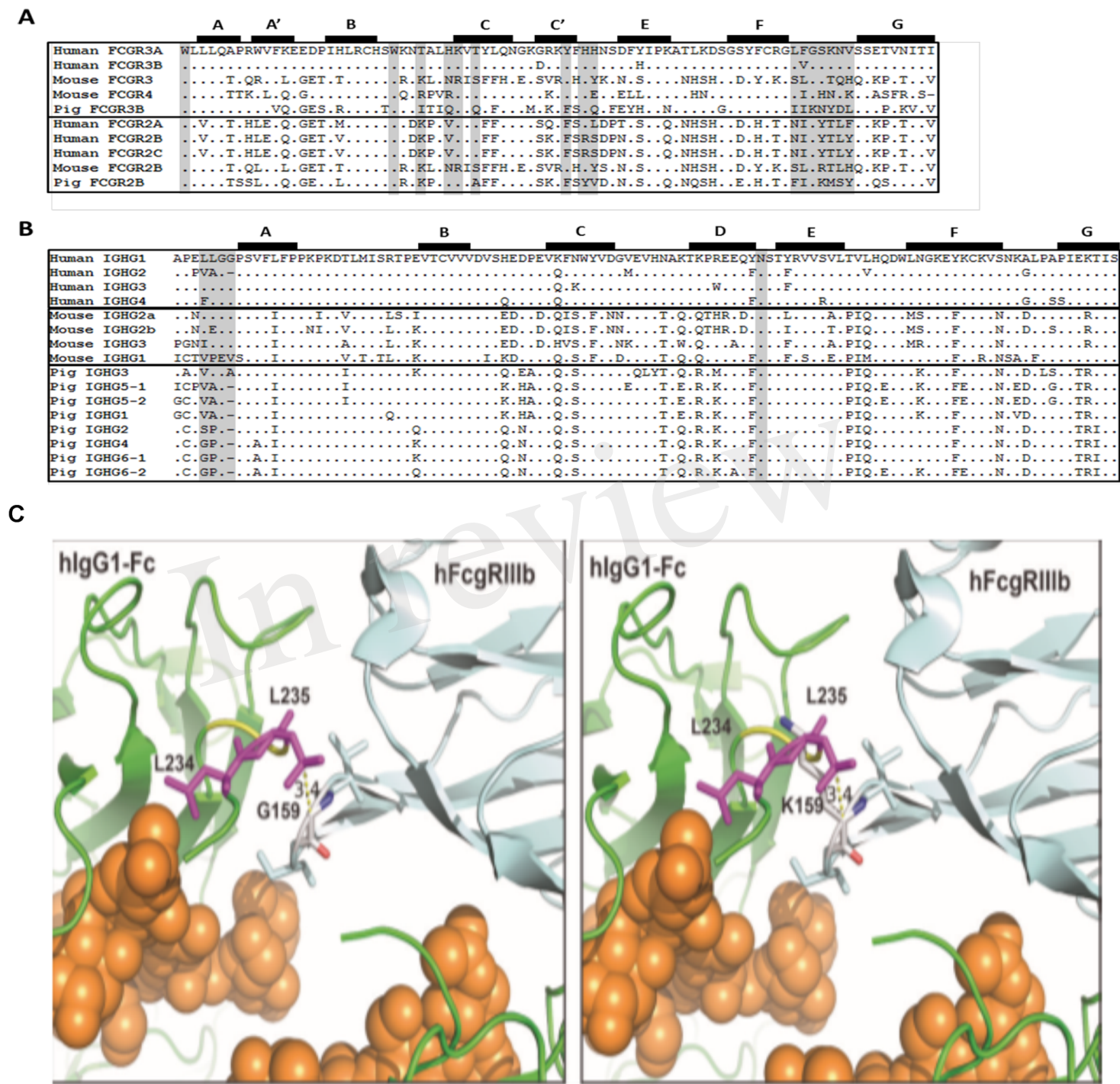


Figure 7